Insights into mechanisms of tumor dissemination from circulating tumor cell lines of small cell lung cancer

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ABSTRACT

Despite the fact that the majority of cancer patients succumb to metastatic disease, most aspects of tumor metastasis are not understood in detail at present. Cell biologic steps of dissemination are difficult to characterize in human tumors and research is in large part confined to cell line and experimental animal studies. Epithelial-mesenchymal transition (EMT), invasation of malignant cells, dissemination as circulating tumor cells (CTCs) and eventually mesenchymal-epithelial transition (MET) at distal sites are steps believed to be involved in metastasis. Small cell lung cancer (SCLC) is distinguished by early dissemination and excessive numbers of CTCs, which allowed for the ex vivo expansion of six permanent CTC lines taken from relapsed patients. Cells exhibit an epithelial phenotype with partial EMT traits and are chemoresistant due to formation of large tumorospheres. Since cells may have invaded without undergoing EMT, the role of MET is uncertain. These SCLC CTC cell lines seem to represent the metastasis-inducing cancer cells; these are the minute subpopulation of CTCs capable of surviving in the circulation and transitioning to metastases, leading in turn to resistance and failure of therapy. Full characterization of these lines is expected to provide the markers to find the relevant CTCs among the highly heterogeneous population observable in the context of tumor recurrence.

Key words: Cancer metastasis, small cell lung cancer, circulating tumor cells, epithelial-mesenchymal transition, tumorosphere, chemotherapy, drug resistance

INTRODUCTION

Early detection, precise diagnosis and monitoring of the course of disease during therapy are the objective of individualized care in current oncology. For most tumors, a tumor biopsy is costly, painful, or potentially hazardous for the patient and is not performed during the further development of the malignancy. Especially in tumor types showing a high frequency of systemic disease, such as small cell lung cancer (SCLC), a small needle biopsy is procured first to confirm the diagnosis, and chemotherapy is started without any further invasive procedure. Thus, the opportunity to obtain essential information from blood samples, as so-called
liquid biopsies, would offer significant advantages. Non-invasive detection and monitoring of patient tumors, employing cell-free circulating tumor DNA (ctDNA) or circulating tumor cells (CTCs), have been employed for investigations into multiple tumor types. CTCs are described as cells, shed by primary or secondary tumors into vasculature, that keep circulating in the bloodstream of cancer patients. Reports indicate that patients with lower counts of CTCs survive longer than the patients with higher CTCs counts. For example, lower numbers of CTCs were observed for 21 patients with limited SCLC (median = 6, range 0-220) compared with 38 patients with extensive stage (median = 63, range 0-14,040) and the absence of measurable CTCs in 27% of patients was correlated with prolonged survival (hazard ratio: 3.4; P ≤ 0.001).

Furthermore, CTC counts have been proposed as a surrogate marker for assessment of responses to therapy in cancer patients to facilitate more rapid drug evaluation. The Food and Drug Administration approved CellSearch® system (Veridex, Raritan, NJ, USA) enumerates intact CTCs for a prognosis of overall survival for breast, prostate and colon cancer. For example, CTC count is a robust independent prognostic factor for progression-free recovery and overall survival in patients with early and metastatic breast cancer. Additionally, CTCs have been detected in patients with chronic obstructive pulmonary disease before the actual occurrence of malign lesions, thus allowing for early diagnosis of lung cancer. However, an advantage of the analysis of ctDNA in liquid biopsy is the detection of molecular changes which are occurring within the tumors in real time, especially during development of drug resistance to targeted agents.

Numerous techniques for the enrichment of CTCs have been developed relying on immunological markers, size, rigidity or dielectric properties. These techniques have been used for genetic characterization, marker analysis and short-term cultures for investigation of their cell biology. Despite a host of studies dealing with CTCs, many questions regarding their generation, shedding, survival in the circulation, chemosensitivity and mechanisms of induction of secondary lesion remain to be fully resolved.

Research on tumor dissemination and CTCs has been hampered by the scarcity or heterogeneity of the enriched cells, as well as the inability to define the characteristics of the actual metastasis-inducing CTCs that are expected to be most relevant for the prognosis of the patients. In this study, we have established for the first time six CTC cell lines from a variety of patients with extended metastatic SCLC and present here the implications of the phenotype of these cell lines for tumor dissemination in this highly aggressive malignancy.

DETECTION AND ENRICHMENT OF CTCs

In most tumor types, CTCs are rare events, with a frequency of approximately one CTC among 1-10 million mononuclear blood cells. Therefore, these cancer cells have to be enriched by various methods for further analysis. The probability to detect CTCs in a limited volume of blood has been reviewed by Gkountela et al. and the reported probability of collecting ≥ 1 CTCs in one aliquot of 7.5 mL blood from a patient with 500 CTCs is 50%. Therefore, 20 mL of whole blood would have to be assessed if the cell event were to be detected for a lower frequency at one CTC in 10⁷ leukocytes. The frequency of the CTC population measured in an aliquot may not be a statistical representative of the entire sample.

There are a number of questions remaining for the detection of CTCs and their relationship to their parent bulk tumors. CTCs seem to stem from the frontier of the tumor and it is generally known that the tumor margin is different from the main tumor mass. Most of the CTCs populations analyzed exhibit heterogeneity, pointing to release of different cell populations from distinct regions of the tumor or of metastatic lesions. CTCs can also form clusters including immune cells in the circulation, or be engulfed by the platelets before eventually adhering to the walls of the capillaries and initiating extravasation.

The role of CTCs as prognostic marker and as possible surrogate indicators for response to therapy is discussed in numerous reviews. CTCs which are shed from tumors are mediators of metastatic dissemination and form micrometastasis at distant organs. In positive selection, surface markers of CTCs are targeted, whereas, in negative selection, depletion of blood cells other than CTCs is achieved by targeting their surface markers. The CellSearch® system is by far the most common system for extraction and enumeration of CTCs for clinical investigations. CTCs which have downregulated EpCAM remain undetected throughout this process. By alternative methods, cell size-based sorting is accomplished using microfluidic technology, as with the Parsortix system (Angle, Guildford, UK). Isolation by Size of Tumor cells (ISET®) is another filter-based established method which is used for such cell size-based sorting.

CTCs derived from breast cancer patients are among the most extensively studied for diagnosis.
and treatment.\textsuperscript{[21]} The presence of CTCs, despite ongoing treatment, has proven to be an indicator of worse overall survival; therefore, in one group of HER2\textsuperscript{+} breast cancer patients, HER2\textsuperscript{-} CTCs were identified and trastuzumab-based therapy applied to these patients.\textsuperscript{[17,22,23]} Most of the CTCs isolated from breast cancer patients show the presence of epithelial-mesenchymal transition (EMT) markers such as ET5V, NOTCH1, SNAIL, TGFB1, ZEB1, and ZEB2.\textsuperscript{[24]} Breast cancer patients who showed remaining CTCs after first cycles of chemotherapy progressed rapidly to metastatic disease.\textsuperscript{[25]} In prostate cancer, CTCs have been proposed to act as intermediate or surrogate endpoints for survival and to shorten timelines for drug approval.\textsuperscript{[20]} Patients with lower levels of CTCs have shown slower disease progression in comparison to those having higher levels of CTCs.\textsuperscript{[27]} In colon cancer, CTCs were found as individual cells or as clusters (CTMs) by a CK-based, immunomagnetic cell separation method.\textsuperscript{[28]} CTMs are of particular interest as they are considered to be markers of increased metastatic potential.\textsuperscript{[29]} In lung cancer, CellSearch\textsuperscript{®} and ISET kits indicated a higher number of CTCs in SCLC than NSCLC, reported to be associated with larger tumor size and bone metastasis.\textsuperscript{[30,31]} The high numbers of CTCs in SCLC allowed for their enrichment and initiation of xenografts which seem to resemble the tumor characteristics of the respective patients.\textsuperscript{[32]} Furthermore, a trial to establish ex vivo expanded CTC cell lines was successful and has resulted in the availability of six lines from relapsed patients so far.\textsuperscript{[33]} Characteristics of the first CTC cell lines and their interaction with macrophages have been published.\textsuperscript{[34]}

**SHELDING OF CTCS FROM TUMORS**

Release of CTCs into the circulation is frequently termed shedding, a designation for a process for which the details are not known. CTCs are reported to be shed from solid tumors at a daily rate of 3.2 to 4.1 \times 10^6 per gram of tissue, based on a single artificial rat model.\textsuperscript{[35]} In one study, the rate of tumor cell shedding into efferent blood was measured in both growing and regressing MTW9 rat mammary carcinomas. Cell shedding rates of growing versus regressing tumors were not significantly different over a tumor size range of 2-4 g. Half of these CTCs perished within 2.4 h, although longer half-lives were reported in a clinical setting.\textsuperscript{[28,36]} Tumor cells are rapidly cleared from circulating blood and a 2-g MTW9 carcinoma reportedly released enough cells into the circulation to transplant the tumor every 24 h, although the majority of the cells were reported to be apoptotic/necrotic.\textsuperscript{[30]} The cell loss via blood comprised about 10% of the tumor weight and resulted in a CTC count of approximately 20,000 CTCs/mL blood. Clearly, this estimation of the release of CTCs reported from an experimental animal model cannot be extrapolated to human tumors. No form of the shedding of CTCs causes a human tumor to lose 10% of its size in one day, nor is a CTC count of 20,000 cells/mL observed in most cancer patients. A threshold of 5 CTCs/7.5 mL blood has been defined by the Cellsearch\textsuperscript{®} system for breast and prostate cancer, and a lower threshold of 3 CTCs/7.5 mL blood has been defined for colon cancer patients. These figures are for favorable or poor prognosis, respectively.\textsuperscript{[7,37]} Consequently, the attrition rate in the circulation based on this artificial animal model seems to be a considerable overestimation. The specific mechanisms of tumor cell shedding are not known at present. CTCs seem to origin as specialized cell types, different from the bulk of the tumor cells, from the borders of the tumor. CTCs leave the particular microenvironmental milieu characterized by inflammation, acidosis and hypoxia through the interaction of a host of participating cell types. Therefore, CTCs are not expected to represent the bulk of tumor cells and are not typical of the cell biologic behavior and chemoresistance of the main body of the tumor. SCLC extended disease responds well to the first cycles of platinum-based chemotherapy but recurs within approximately one year as tumors which exhibit universal chemoradioresistance.\textsuperscript{[38]} Contrary to expectations, the first two SCLC CTC cell lines proved to be chemosensitive to the second-line chemotherapeutics topotecan and epirubicin although some tumor cells must have survived the initial successful treatment and eventually give rise to chemoresistant relapses.\textsuperscript{[39]} Therefore, the use of CTCs as surrogate markers for the bulk tumor is questionable.

**EMT IN TUMOR CELL SHEDDING**

A general assumption supposes that tumor cells invade through a process termed EMT.\textsuperscript{[40]} Accordingly, epithelial tumor cell downregulate epithelial markers, such as E-cadherin, EpCAM and cytokeratins, eventually expressing mesenchymal markers such as vimentin, neural cell adhesion molecule (NCAM) and others. In this way, cells gain mobility and migrate to intravasate and reach distant sites to establish secondary lesions. EMT is regulated by a number of specific transcription factors belonging to the SNAIL, TWIST and ZEB families and is modulated by microenvironmental conditions, inflammatory cytokines and chemotherapy.\textsuperscript{[41]} Since it has proven difficult to demonstrate tumor cells with EMT traits in patients, incomplete EMT or transitional EMT has been proposed as a model. This model
presupposes no complete switch in phenotype, but rather a type of transition which may be as minimal as a slight downregulation of epithelial features. EMT phenotypes have been reported among heterogeneous CTC populations and increased fractions of cells with such mesenchymal features have been demonstrated to correlate with a poorer prognosis in breast cancer patients.[24]

It may still be possible that tumor cells with epithelial characteristics enter the bloodstream without undergoing EMT. An alternative model, termed “cooperative migration”, posits that EMT-type cells help epithelial cells to gain access to the circulation, but to reside at the bulk tumor. Excessive numbers of CTCs have been observed in SCLC, which is frequently associated with local inflammation and in inflammatory breast cancer. Thus, immune cells and inflammation may promote release of tumor cells into the circulation possibly without EMT. The SCLC CTC cell lines have been found to recruit macrophages and to lack a phenotypic switch with full expression of mesenchymal traits.[34,42] In conclusion, complete or partial EMT is not proven to be a prerequisite to disseminate tumor cells, and therapeutic options to inhibit such a transition need to be considered cautiously.[43]

SURVIVAL CTCS IN THE CIRCULATION

The great majority of CTCs seem to be short-lived and to perish in the circulation. Of the several forms of CTCs shed by the primary tumor, only about 0.1% survive in the circulation and only about 0.01% is responsible for metastasis.[8,10] This attrition has been attributed to shear stress and an unfavorable microenvironment too different from the local tumor conditions. CTCs in the hematogenous circulation must survive a variety of stresses, and epithelial cells may undergo anoikis in the absence of cellular attachment.[44] The vast majority of CTCs are likely to become trapped in various capillary beds. They are destroyed by hemodynamic shear forces and predation by cells of the innate immune system – specifically natural killer cells. Consistent with this view, a great deal of CTC-associated material is detectable in CTC-positive tumor patients.[15] Reported half-lives have ranged from several hours, according to experimental animal models, to long-term persistence. The SCLC CTC lines show continuing disposal of microparticles and cellular fragments, thus generating CTC associated materials under optimal conditions in tissue culture in the absence of shear stress.[42] Partial disintegration of CTC tumorospheres may function as a source of decoy material to protect other CTCs and the bulk tumor from attacks by both the immune system and chemotherapeutics.

INDUCTION OF EXTRAVASATION AND MESENCHYMAL-EPITHELIAL TRANSITION

In general, the secondary lesions induced by CTCs show an epithelial phenotype similar to the originating primary tumor.[40] Provided that the metastases were established by cells which underwent EMT, at some point during tumor spread and extravasation this transition has to be reversed through a process termed mesenchymal-epithelial transition (MET). This process has not been observed directly, but has been inferred from the mesenchymal traits of disseminated/CTCs and the histology of secondary lesions.[46] The factors causing this supposedly phenotypic switch, along with their possible derivation from the cancer cells themselves (seed) or the metastatic site (soil) are largely unknown. In the case of SCLC CTC lines, the cells are positive for EpCAM, E-cadherin and proteins involved in cell junctions and form spontaneously typical large spheroids with diameters of up to 1-2 mm in regular tissue culture medium without any factors preventing adhesion to cell culture flasks.[43] Although expression of vimentin and NCAM is observed, the formation of these organized and large spheres is a typical epithelial feature not observed in SCLC tumor cell lines in vitro. Since these tumorospheres develop from CTCs of relapsed SCLC patients within a short time, and are found in cell suspension derived from xenografts induced by such CTCs, they seem not to stem from an in vitro transition in tissue culture but to present the original in vivo phenotype. Thus, these metastasis-inducing CTCs seem to be present as cancer cells exhibiting an epithelial phenotype and organization and may be trapped in capillaries or reside in protected sites, possibly in a dormant state. Tumorospheres exceeding diameters of 2 mm during their development tend to disintegrate and may be the source of non-proliferating cell clusters observed in the blood of metastatic cancer patients.[10]

METASTASIS AND DRUG RESISTANCE

Metastases not only damage secondary organs and exacerbate the deleterious effects of malignancies in general but frequently exhibit chemoresistance to reinitiation of primary or second-line chemotherapy agents. Especially in SCLC, excellent response rates to initial chemotherapy can be followed by relapses within approximately one year, which exhibit broad chemoresistance and result in failure of treatment.[1,38] Attempts have been made to characterize the chemosensitivity of CTCs in short term cultures in various tumor entities.[46] However, CTCs are specialized cells different from the tumor bulk and most likely also
from the developing metastases. It was universally assumed that CTCs survive as chemoresistant cells and may predict the responsiveness of resulting secondary lesions. Surprisingly, the SCLC CTC lines were found to be highly chemosensitive to agents of common use in this tumor entity, except the CTC line established from a patient refractory to initial therapy with cisplatin.\(^{39}\) Thus, some tumor cells have survived the initial cycles of chemotherapy, possibly in an inflammatory environment, and seem to develop a kind of chemoresistance not accomplished at the cellular level.\(^{34}\) Tumorospheres are known to be comprised of a small region of proliferating cells and layers of quiescent cells surrounding an inner hypoxic core which provides protection against irradiation due to the lack of reactive oxygen species [Figure 1].\(^{47,48}\)

Therapy of SCLC has not been improved for the last several decades despite the clinical testing of a host of chemotherapeutics covering the widest range of clinical targets available.\(^{1,49}\) More than 600 trials exploring therapeutic interventions in SCLC are currently in the U.S. clinical trials registry, National Institutes of Health.\(^{50}\) Since it is not reasonable to assume that SCLC tumors express a host of individual molecular mechanisms, tumorospheres provide an alternative explanation for chemoresistance, in which potential chemotherapeutic drugs are prevented from reaching their cellular target (in responsive cells) in sufficient quantities.\(^{51}\) Whether the validity of this kind of chemoresistance, in the form of tumor spheroids, is confined to SCLC or is common to other tumor types, remains to be investigated.

**CONCLUSION**

Mechanisms of tumor metastasis have been investigated using numerous cell culture studies and research employing experimental animal models as well as analyses of clinical specimens. Models have been proposed for the discrete steps of tumor metastasis, but the actual dissemination of malignancies in patients is difficult to assess. Clearly, CTCs are instrumental in translocation of tumor cells to distal sites and in the induction of secondary lesions. Although CTC counts have shown prognostic significance, their limited accessibility and marked heterogeneity have limited their usefulness.

A panel of permanent CTC lines from SCLC patients revealed absence of full EMT, presence of chemosensitivity, and an epithelial phenotype with formation of tumorospheres as physical barrier against chemoradiotherapy. Unfortunately, there are currently no means available to target these spheroid structures in a clinical setting, and further investigation is needed to study the cell biology of CTC aggregates in detail, and to study overcoming resistance using targeted agents involving enzymes, cell junctions opener, nanomaterials and other mechanisms.

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**Conflicts of interest**

There are no conflicts of interest.

**Patient consent**

Informed patient consent was obtained throughout the study.

**Ethics approval**

The respective ethics protocol (Approval of the Medical University of Vienna Nr. 366/2003 and amendments obtained prior to the commencement of the study) was followed throughout the study.

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